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# (54) Dye-donor element containing transferable protection overcoat

(57) A dye-donor element for thermal dye transfer comprising a support having thereon at least one dye layer area comprising an image dye in a binder and another area comprising a transferable protection layer, the transferable protection layer area being approximately equal in size to the dye layer area, wherein the transferable protection layer contains inorganic particles, a polymeric binder and unexpanded synthetic thermoplastic polymeric microspheres, the microspheres having a particle size in the unexpanded condition of from 5 to 20 μm, and which expand to 20 to 120 μm upon

application of heat during transfer of the protection layer to an image-receiving layer to provide a matte surface thereon, the microspheres comprising a mixture of low softening point microspheres and high softening point microspheres, the low softening point microspheres having a softening point less than 105° C, the high softening point microspheres having softening point greater than 110°C, and the ratio of the low softening point microspheres to the high softening point microspheres being from 9:1 to 1:6.

#### Description

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[0001] This invention relates to a dye-donor element for thermal dye transfer, and more particularly to the use of a transferable protection overcoat in the element for transfer to a thermal print to provide a matte surface thereon.

[0002] In recent years, thermal transfer systems have been developed to obtain prints from pictures that have been generated electronically from a color video camera. According to one way of obtaining such prints, an electronic picture is first subjected to color separation by color filters. The respective color-separated images are then converted into electrical signals. These signals are then operated on to produce cyan, magenta and yellow signals. These signals are then transmitted to a thermal printer. To obtain the print, a cyan, magenta or yellow dye-donor element is placed face-to-face with a dye-receiving element. The two are then inserted between a thermal printing head and a platen roller. A line-type thermal printing head is used to apply heat from the back of the dye-donor sheet. The thermal printing head has many heating elements and is heated up sequentially in response to one of the cyan, magenta and yellow signals. The process is then repeated for the other two colors. A color hard copy is thus obtained which corresponds to the original picture viewed on a screen. Further details of this process and an apparatus for carrying it out are contained in U.S. Patent 4,621,271.

[0003] Thermal prints are susceptible to retransfer of dyes to adjacent surfaces and to discoloration by fingerprints. This is due to dye being at the surface of the dye-receiving layer of the print. These dyes can be driven further into the dye-receiving layer by thermally fusing the print with either hot rollers or a thermal head. This will help to reduce dye retransfer and fingerprint susceptibility, but does not eliminate these problems. However, the application of a protection overcoat will practically eliminate these problems. This protection overcoat is applied to the receiver element by heating in a likewise manner after the dyes have been transferred. The protection overcoat will improve the stability of the image to light fade and oil from fingerprints.

[0004] In a thermal dye transfer printing process, it is desirable for the finished prints to compare favorably with color photographic prints in terms of image quality. The look of the final print is very dependent on the surface texture and gloss. Typically, color photographic prints are available in surface finishes ranging from very smooth, high gloss to rough, low gloss matte.

[0005] If a matte finish is desired on a thermal print, it has been previously accomplished by using matte sprays or by matte surface applications through post printing processors. However, both of these solutions are costly and add a degree of complexity to the process.

[0006] Serial No. 09/550,367 of Simpson et al., filed April 19, 2000, and JP 09/323482 relate to the use of expandable microspheres in a transferable protection layer area of a dye-donor element. However, there is a problem with these microspheres in that they will not provide a defect-free print with a desired gloss at a low printhead temperature.

[0007] It is the object of this invention to provide a dye-donor element for thermal dye transfer printing that can impart a matter or low gloss finish onto a receiving element. It is another object of this invention to provide a dye-donor element for thermal dye transfer printing that provides a protection layer which improves the adhesion between the protection layer and the receiving layer resulting in less defects. It is another object of the invention to provide a dye-donor element for thermal dye transfer printing that provides a protection layer wherein the gloss can be varied.

[0008] These and other objects are achieved in accordance with this invention which relates to a dye-donor element for thermal dye transfer comprising a support having thereon at least one dye layer area comprising an image dye in a binder and another area comprising a transferable protection layer, the transferable protection layer area being approximately equal in size to the dye layer area, wherein the transferable protection layer contains inorganic particles, a polymeric binder and unexpanded synthetic thermoplastic polymeric microspheres, the microspheres having a particle size in the unexpanded condition of from 5 to 20 µm, and which expand to 20 to 120 µm upon application of heat during transfer of the protection layer to an image-receiving layer to provide a matte surface thereon, the microspheres comprising a mixture of low softening point microspheres and high softening point microspheres, the low softening point microspheres having a softening point less than 105°C, the high softening point microspheres having softening point greater than 110°C, and the ratio of the low softening point microspheres to the high softening point microspheres being from 9:1 to 1:6, preferably from 4:1 to 1:4.

[0009] By use of the invention, a dye-donor element is provided containing a transferable protection layer which is capable of giving a variable gloss.

[0010] During application of the protection layer to the receiver element, heat from the linear thermal printing head causes the microspheres to expand to many times their original size. This causes a roughening of the surface to occur resulting in a matte or lower gloss image comparable to that obtained on a matte surface photographic paper. In accordance with the invention, a mixture of microspheres that differ in the softening temperature, or T-start of the microsphere wall is used. When a low printhead temperature is employed, a 60° gloss value of 65 or greater is obtained, while at a high print head temperature, a 60° gloss value of less than 40 is obtained. Thus, by merely varying the temperature of transfer of the protection layer, any desired degree of gloss may be obtained. The temperature of transfer of the protection layer can be varied by changing the power supplied to the thermal print head.

[0011] In a preferred embodiment of the invention, the dye-donor element is a multicolor element comprising repeating color patches of yellow, magenta and cyan image dyes, respectively, dispersed in a binder, and a patch containing the protection layer.

[0012] In another embodiment of the invention, the protection layer is the only layer on the donor element and is used in conjunction with another dye-donor element which contains the image dyes.

[0013] In another preferred embodiment of the invention, the dye-donor element is a monochrome element and comprises repeating units of two areas, the first area comprising a layer of one image dye dispersed in a binder, and the second area comprising the protection layer.

[0014] In another preferred embodiment of the invention, the dye-donor element is a black-and-white element and comprises repeating units of two areas, the first area comprising a layer of a mixture of image dyes dispersed in a binder to produce a neutral color, and the second area comprising the protection layer.

[0015] Any expandable microspheres may be used in the invention provided they have the softening point parameters as described above. Materials which can be used are disclosed, for example, in U.S. Patents 3,556,934 and 3,779,951.

[0016] In a preferred embodiment of the invention, the expandable microspheres are white, spherically-formed, hollow particles of a thermoplastic shell encapsulating a low-boiling, vaporizable substance, such as a liquid, which acts as a blowing agent. When the unexpanded microspheres are heated, the thermoplastic shell softens and the encapsulated blowing agent expands, building pressure. This results in expansion of the microsphere. The various expandable microspheres differ in the temperature where the microspheres began to expand. This is known as the softening point, or T-start temperature.

[0017] The expandable microspheres employed in the invention may be formed by encapsulating isopentane, isobutane or any other low-boiling, vaporizable substance into a microcapsule of a thermoplastic resin such as a vinylidene chloride-acrylonitrile copolymer, a methacrylic acid ester-acrylonitrile copolymer or a vinylidene chloride-acrylic acid ester copolymer. These microspheres are available commercially as Expancel ® Microspheres 461-20-DU, 6-9 µm particle diameter weighted average, T-start 98-104 ° C, (Expancel Inc.); Expancel ® Microspheres 461-DU, 9-15 µm particle diameter weighted average, T-start 98-104 °C, (Expancel Inc.); and Expancel ® Microspheres 091-DU, 10-16 µm particle diameter weighted average, T-start 118-126 °C, (Expancel Inc.). In a preferred embodiment of the invention, the low softening point microspheres have a softening point from 98 °C to 104 °C. In another preferred embodiment of the invention, the high softening point microspheres have a softening point from 118 °C to 126 °C.

[0018] The present invention provides a protection overcoat layer on a thermal print by uniform application of heat using a thermal head. After transfer to the thermal print, the protection layer provides superior protection against image deterioration due to exposure to light, common chemicals, such as grease and oil from fingerprints, and plasticizers from film album pages or sleeves made of poly(vinyl chloride). The protection layer is generally applied at a coverage of at least 0.03 g/m $^2$  to 1.5 g/m $^2$  to obtain a dried layer of less than 1  $\mu$ m.

[0019] As noted above, the transferable protection layer comprises the microspheres dispersed in a polymeric binder. Many such polymeric binders have been previously disclosed for use in protection layers. Examples of such binders include those materials disclosed in U.S. Patent 5,332,713. In a preferred embodiment of the invention, poly(vinyl acetal) is employed.

[0020] Inorganic particles are present in the protection layer of the invention. There may be used, for example, silica, titania, alumina, antimony oxide, clays, calcium carbonate, talc, etc. as disclosed in U.S. Patent 5,387,573. In a preferred embodiment of the invention, the inorganic particles are silica. The inorganic particles improve the separation of the laminated part of the protection layer from the unlaminated part upon printing.

[0021] In a preferred embodiment of the invention, the protection layer contains from 5% to 60% by weight inorganic particles, from 25% to 60% by weight polymeric binder and from 5% to 60% by weight of the unexpanded synthetic thermoplastic polymeric microspheres.

[0022] In use, yellow, magenta and cyan dyes are thermally transferred from a dye-donor element to form an image on the dye-receiving sheet. The thermal head is then used to transfer the clear protection layer, from another clear patch on the dye-donor element or from a separate donor element, onto the imaged receiving sheet by uniform application of heat. The clear protection layer adheres to the print and is released from the donor support in the area where heat is applied.

[0023] Any dye can be used in the dye layer of the dye-donor element of the invention provided it is transferable to the dye-receiving layer by the action of heat. Especially good results have been obtained with sublimable dyes. Examples of sublimable dyes include anthraquinone dyes, e.g., Sumikaron Violet RS® (Sumitomo Chemical Co., Ltd.), Dianix Fast Violet 3R FS® (Mitsubishi Chemical Industries, Ltd.), and Kayalon Polyol Brilliant Blue N BGM® and KST Black 146® (Nippon Kayaku Co., Ltd.); azo dyes such as Kayalon Polyol Brilliant Blue BM®, Kayalon Polyol Dark Blue 2BM®, and KST Black KR® (Nippon Kayaku Co., Ltd.), Sumikaron Diazo Black 5G® (Sumitomo Chemical Co., Ltd.), and Miktazol Black 5GH® (Mitsui Toatsu Chemicals, Inc.); direct dyes such as Direct Dark Green B® (Mitsubishi Chemical Industries, Ltd.) and Direct Brown M® and Direct Fast Black D® (Nippon Kayaku Co. Ltd.); acid dyes such as Kayanol Milling Cyanine 5R® (Nippon Kayaku Co. Ltd.); basic dyes such as Sumiacryl Blue 6G® (Sumitomo Chemical

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Co., Ltd.), and Aizen Malachite Green® (Hodogaya Chemical Co., Ltd.);

$$CH_{3}$$

$$CH_{3}$$

$$CH - CH = \begin{cases} N - C_{6}H_{5} \\ N \end{cases} \quad (yellow)$$

$$C_{2}H_{5}$$

$$N(CH_{3})_{2}$$

$$(C_2H_5)_2N \longrightarrow CH \longrightarrow N \longrightarrow N$$

$$(Yellow)$$

$$N(CH_3)_2$$

CONHCH<sub>3</sub> (cyan)
$$N \longrightarrow N(C_2H_5)_2$$

CONHCH<sub>3</sub> (cyan)
$$N \longrightarrow N (C_2H_5)_2$$

or any of the dyes disclosed in U.S. Patent 4,541,830. The above dyes may be employed singly or in combination to obtain a monochrome. The dyes may be used at a coverage of from 0.05 to 1 g/m² and are preferably hydrophobic. [0024] A dye-barrier layer may be employed in the dye-donor elements of the invention to improve the density of the

transferred dye. Such dye-barrier layer materials include hydrophilic materials such as those described and claimed in U.S. Patent 4.716.144.

[0025] The dye layers and protection layer of the dye-donor element may be coated on the support or printed thereon by a printing technique such as a gravure process.

[0026] A slipping layer may be used on the back side of the dye-donor element of the invention to prevent the printing head from sticking to the dye-donor element. Such a slipping layer would comprise either a solid or liquid lubricating material or mixtures thereof, with or without a polymeric binder or a surface-active agent. Preferred lubricating materials include oils or semi-crystalline organic solids that melt below 100°C such as poly(vinyl stearate), beeswax, perfluorinated alkyl ester polyethers, poly-caprolactone, silicone oil, poly(tetrafluoroethylene), carbowax, poly(ethylene glycols), or any of those materials disclosed in U.S. Patents 4,717,711; 4,717,712; 4,737,485; and 4,738,950. Suitable polymeric binders for the slipping layer include poly(vinyl alcohol-co-butyral), poly(vinyl alcohol-co-acetal), polystyrene, poly(vinyl acetate), cellulose acetate butyrate, cellulose acetate propionate, cellulose acetate or ethyl cellulose.

[0027] The amount of the lubricating material to be used in the slipping layer depends largely on the type of lubricating material, but is generally in the range of 0.001 to 2 g/m². If a polymeric binder is employed, the lubricating material is present in the range of 0.05 to 50 weight %, preferably 0.5 to 40 weight %, of the polymeric binder employed.

[0028] Any material can be used as the support for the dye-donor element of the invention provided it is dimensionally stable and can withstand the heat of the thermal printing heads. Such materials include polyesters such as poly(ethylene terephthalate); polyamides; polycarbonates; glassine paper; condenser paper; cellulose esters such as cellulose acetate; fluorine polymers such as poly(vinylidene fluoride) or poly(tetrafluoroethylene-co-hexafluoropropylene), polyethers such as polyoxymethylene; polyacetals; polyolefins such as polystyrene, polyethylene, polypropylene or methylpentene polymers; and polyimides such as polyimide amides and polyetherimides. The support generally has a thickness of from 2 to 30  $\mu$ m.

[0029] The dye-receiving element that is used with the dye-donor element of the invention usually comprises a support having thereon a dye image receiving layer. The support may be a transparent film such as a poly(ether sulfone), a polyimide, a cellulose ester such as cellulose acetate, a poly(vinyl alcohol-co-acetal) or a poly(ethylene terephthalate). The support for the dye-receiving element may also be reflective such as baryta-coated paper, polyethylene-coated paper, white polyester (polyester with white pigment incorporated therein), an ivory paper, a condenser paper or a synthetic paper such as DuPont Tyvek®.

[0030] The dye image-receiving layer may comprise, for example, a polycarbonate, a polyurethane, a polyester, poly (vinyl chloride), poly(styrene-co-acrylonitrile), polycaprolactone or mixtures thereof. The dye image-receiving layer may be present in any amount which is effective for the intended purpose. In general, good results have been obtained at a concentration of from 1 to 5 g/m<sup>2</sup>.

[0031] As noted above, the dye donor elements of the invention are used to form a dye transfer image. Such a process comprises imagewise heating a dye-donor element as described above and transferring a dye image to a dye receiving element to form the dye transfer image. After the dye image is transferred, the protection layer is then transferred on top of the dye image.

[0032] The dye donor element of the invention may be used in sheet form or in a continuous roll or ribbon. If a continuous roll or ribbon is employed, it may have only one dye or may have alternating areas of other different dyes, such as sublimable cyan and/or magenta and/or yellow and/or black or other dyes. Such dyes are disclosed in U.S. Patents 4,541,830; 4,698,651; 4,695,287; 4,701,439; 4,757,046; 4,743,582; 4,769,360 and 4,753,922. Thus, one-, two-, three- or four-color elements (or higher numbers also) are included within the scope of the invention.

[0033] In a preferred embodiment of the invention, the dye-donor element comprises a poly(ethylene terephthalate) support coated with sequential repeating areas of yellow, cyan and magenta dye, and the protection layer noted above, and the above process steps are sequentially performed for each color to obtain a three-color dye transfer image with a protection layer on top. Of course, when the process is only performed for a single color, then a monochrome dye transfer image is obtained.

[0034] Thermal printing heads which can be used to transfer dye from the dye-donor elements of the invention are available commercially. There can be employed, for example, a Fujitsu Thermal Head FTP-040 MCSOO1, a TDK Thermal Head LV5416 or a Rohm Thermal Head KE 2008-F3.

[0035] A thermal dye transfer assemblage of the invention comprises

- (a) a dye-donor element as described above, and
- (b) a dye-receiving element as described above,

the dye receiving element being in a superposed relationship with the dye donor element so that the dye layer of the donor element is in contact with the dye image-receiving layer of the receiving element.

[0036] The above assemblage comprising these two elements may be preassembled as an integral unit when a monochrome image is to be obtained. This may be done by temporarily adhering the two elements together at their

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margins. After transfer, the dye-receiving element is then peeled apart to reveal the dye transfer image.

[0037] When a three-color image is to be obtained, the above assemblage is formed on three occasions during the time when heat is applied by the thermal printing head. After the first dye is transferred, the elements are peeled apart. A second dye-donor element (or another area of the donor element with a different dye area) is then brought in register with the dye-receiving element and the process is repeated. The third color is obtained in the same manner. Finally, the protection layer is applied on top.

[0038] The following example is provided to illustrate the invention.

# Control Element C-1 (Mixture of Microspheres With Same Softening Point)

[0039] Protection layer donor elements were prepared by coating on the back side of a 6  $\mu$ m poly(ethylene terephthalate) support:

1) a subbing layer of titanium alkoxide, Tyzor TBT®, (DuPont Corp.) (0.13 g/m²) from a n-propyl acetate and n-butyl alcohol solvent mixture (85/15), and

2) a slipping layer containing an aminopropyl-dimethyl-terminated polydimethylsiloxane, PS513® (United Chemical Technologies) (0.01 g/m²), a poly(vinyl acetal) binder, KS-1, (Sekisui Co.), (0.38 g/m²), p-toluenesulfonic acid (0.0003 g/m²) and candellila wax (0.02 g/m²) coated from a solvent mixture of diethylketone, methanol and distilled water (88.7/9.0/2.3).

[0040] On the front side of the element was coated a transferable overcoat layer of poly(vinyl acetal), KS-1, (Sekisui Co.), at a laydown of 0.432 g/m², colloidal silica, MA-ST-M (Nissan Chemical Co.), at a laydown of 0.335 g/m², Expancel® microspheres 461-20-DU (Expancel Inc.), (softening point of 98-104°C) at a laydown of 0.099 g/m², and Expancel® microspheres 461-DU (Expancel Inc.), (softening point of 98-104°C) at a laydown of 0.149 g/m², coated from a 75% 3-pentanone and 25% methanol solvent mixture.

## Control Element C-2

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[0041] This element is the same as C-1 with the addition to the transferable protection overcoat layer of poly(vinyl butyral) Butvar B-76®, (Solutia Inc.) at a laydown of 0.043 g/m².

## Control Element C-3

[0042] This element is the same as C-2 except that the colloidal silica was coated at a laydown of 0.439 g/m², the 461-20-DU microspheres were coated at a laydown of 0.164 g/m², and the 461-DU microspheres were coated at a laydown of 0.084 g/m².

## Control Element C-4

[0043] This element is the same as C-1 except that the 461-20-DU microspheres were coated at a laydown of 0.112 g/m², and the 461-DU microspheres were coated at a laydown of 0.168 g/m².

## Control Element C-5

[0044] This element is the same as C-4 with the addition to the transferable protection overcoat layer of poly(vinyl butyral) Butvar B-76®, (Solutia Inc.) at a laydown of 0.043 g/m².

# Element 1 of the Invention (Mixture of Microspheres With Different Softening Points)

[0045] This element was prepared the same as Control Element C-1, except that the Expancel® microspheres 461-DU were replaced with Expancel® microspheres 091-DU (softening point of 118-126°C.)

## Element 2 of the Invention

55 [0046] This element is the same as Element 1 of the Invention with the addition to the transferable protection overcoat layer of poly(vinyl butyral) Butvar B-76®, (Solutia Inc.) at a laydown of 0.043 g/m².

#### Element 3 of the Invention

[0047] This element is the same as Element 2 of the Invention except that the colloidal silica was coated at a laydown of 0.439 g/m $^2$ , the 461-20-DU microspheres were coated at a laydown of 0.164 g/m $^2$ , and the 091-DU microspheres were coated at a laydown of 0.084 g/m $^2$ .

## Element 4 of the Invention

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[0048] This element is the same as Element 1 of the Invention except that the 461-20-DU microspheres were coated at a laydown of 0.112 g/m², and the 091-DU microspheres were coated at a laydown of 0.168 g/m².

#### Element 5 of the Invention

[0049] This element is the same as Element 4 of the Invention with the addition to the transferable protection overcoat layer of poly(vinyl butyral) Butvar B-76®, (Solutia Inc.) at a laydown of 0.043 g/m².

# Receiving Element

[0050] A thermal dye-transfer receiving element was prepared by coating the following layers in order onto a support of an OPPalyte® polypropylene

a) a subbing layer of Prosil® 221 (aminopropyl-triethoxysilane) and Prosil® 2210 (aminofunctional epoxysilane) (PCR, Inc.) (1:1 weight ratio) and LiCl (0.0022 g/m²) in an ethanol-methanol-water solvent mixture. The resultant solution (0.10 g/m²) contained approximately 1% of silane component, 3% water, and 96% of 3A alcohol;

b) a dye-receiving layer containing Makrolon® KL3-1013 (a polyether-modified bisphenol-A polycarbonate block copolymer (BayerAG) (1.52 g/m²), Lexan® 141-112 bisphenol-A polycarbonate (General Electric Co.) (1.24 g/m²), Fluorad® FC-431 a perfluorinated alkylsulfonamidoalkylester surfactant (3M Co.) (0.011g/m²), Drapex® 429 polyester plasticizer (Witco Corp.) (0.23 g/m²), 8 μm crosslinked poly(styrene-co-butyl acrylate-co-divinylbenzene) elastomeric beads (Eastman Kodak Co.) (0.006 g/m²) and diphenyl phthalate (0.46 g/m²) coated from dichloromethane; and

c) a dye-receiver overcoat coated from a solvent mixture of methylene chloride and trichloroethylene containing a polycarbonate random terpolymer of bisphenol-A (50 mole-%), diethylene glycol (49 mole-%), and polydimethylsiloxane (1 mole%) (2,500 MW) block units (0.55 g/m²); a bisphenol A polycarbonate modified with 50 mole-% diethylene glycol (2,000 MW) (0.11 g/m²); Fluorad® FC-431 surfactant (0.022 g/m²); and DC-510® surfactant (Dow Corning Corp.) (0.003 g/m²).

## Polycarbonates used:

## [0051]

 $\begin{array}{c|c}
C H_3 & O \\
C & | \\
C H_2 C H_2 O \\
\hline
C H_3 & n = 1 2 O
\end{array}$ 

KL3-1013, block copolymer of polyether glycol and bisphenol A polycarbonate (Bayer AG)

Bisphenol A polycarbonate Lexan 141® (General Electric Company)

## **Printing**

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[0052] Using Kodak Professional EKTATHERM XLS XTRALIFE Color Ribbon (Eastman Kodak Co. Catalog No. 807-6135) and a Kodak Model 8650 Thermal Printer, a Status A neutral density image with a maximum density of at least 2.3 was printed on the receiver described above. The color ribbon-receiver assemblage was positioned on an 18mm platen roller and a TDK thermal head (No. 3K0345) with a head load of 6.35Kg was pressed against the platen roller. The TDK 3K0345 thermal print head has 2560 independently addressable heaters with a resolution of 300 dots/ inch and an average resistance of 3314 $\Omega$ . The imaging electronics were activated when an initial print head temperature of 36.4°C had been reached. The assemblage was drawn between the printing head and platen roller at 16.9 mm/sec. Coincidentally, the resistive elements in the thermal print head were pulsed on for 58 µsec every 76 µsec. Printing maximum density required 64 pulses "on" time per printed line of 5.0 msec. The voltage supplied was 13.6 volts resulting in an instantaneous peak power of approximately 58.18 x 10-3 Watt/dot and the maximum total energy required to print Dmax was 0.216 mJoules/dot. The process is repeated sequentially, yellow, magenta, cyan to obtain the desired neutral image.

[0053] Each of the protective layer elements described above was placed in contact with the polymeric receiving layer side of the receiver element containing the neutral density image described above. The printing process was used to heat the transferable protection overcoat uniformly with the thermal head to permanently adhere the transferable protection overcoat to the print. The print energy was varied by changing the head voltage and line enable width. The donor support was peeled away as the printer advanced through its heating cycle, leaving the transferable protection overcoat adhered to the imaged receiver.

## Gloss

[0054] The 60° gloss values were measured using a Byk-Gardner Tri-gloss meter. The readings were done with the meter perpendicular to the printing direction and each value is the average of four readings randomly selected on the sample. The following results were obtained:

30 Table

Iabit		
Transferable Protection Overcoat Element	Print Energy (mJ/pixel)	60 ° Gloss
C-1	0.179	64
C-1	0.258	34
C-2	0.179	63
C-2	0.258	30
C-3	0.179	61
C-3	0.258	38
C-4	0.191	61
C-4	0.258	34
C-5	0.191	58
C-5	0.258	31
. 1	0.179	73
1	0.258	36
2	0.179	72
2	0.258	31
3	0.179	66
3	0.258	35
4	0.191	72
4	0.258	34

## Table (continued)

Transferable Protection Overcoat Element	Print Energy (mJ/pixel)	60 ° Gloss
5	0.191	69
5	. 0.258	32

[0055] The above results show that higher gloss can be obtained using the elements of the invention as compared to the control elements at low energy levels. At higher energy levels, the gloss value decreases to give a more matte appearance. In addition, the results also show that different gloss values can be obtained by changing the energy supplied to the thermal print head.

#### Claims

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- 1. A dye-donor element for thermal dye transfer comprising a support having thereon at least one dye layer area comprising an image dye in a binder and another area comprising a transferable protection layer, said transferable protection layer area being approximately equal in size to said dye layer area, wherein said transferable protection layer contains inorganic particles, a polymeric binder and unexpanded synthetic thermoplastic polymeric microspheres, said microspheres having a particle size in the unexpanded condition of from 5 to 20 μm, and which expand to 20 to 120 μm upon application of heat during transfer of said protection layer to an image-receiving layer to provide a matte surface thereon, said microspheres comprising a mixture of low softening point microspheres and high softening point microspheres, said low softening point microspheres having a softening point less than 105°C, said high softening point microspheres having softening point greater than 110°C, and the ratio of said low softening point microspheres to said high softening point microspheres being from 9:1 to 1:6.
- 2. The element of Claim 1 wherein said low softening point microspheres have a softening point from 98 ° C to 104°C.
- 3. The element of Claim 1 wherein said high softening point microspheres have a softening point from 118°C to 126°C.
- 4. The element of Claim 1 wherein said ratio of said low softening point microspheres to said high softening point microspheres being from 4:1 to 1:4.
- 5. The element of Claim 1 wherein said microspheres comprise a vinylidene chloride-acrylonitrile copolymer, a methacrylic acid ester-acrylonitrile copolymer, or a vinylidene chloride-acrylic acid ester copolymer.
- 6. The element of Claim 1 wherein said microspheres comprise an outer shell of a vinylidene chloride-acrylonitrile copolymer, a methacrylic acid ester-acrylonitrile copolymer or a vinylidene chloride-acrylic acid ester copolymer, and a core of a low boiling, vaporizable substance.
- 7. The element of Claim 1 wherein said inorganic particles comprise silica and said polymeric binder is poly(vinyl acetal).
- 8. The element of Claim 1 wherein said protection layer contains from 5% to 60% by weight inorganic particles, from 25% to 60% by weight polymeric binder and from 5% to 60% by weight of unexpanded synthetic thermoplastic polymeric microspheres.
  - 9. A process of forming a protection layer on top of a thermal dye transfer image comprising:
    - (a) imagewise-heating a dye-donor element comprising a support having thereon a dye layer comprising an image dye in a binder, said dye-donor being in contact with a dye-receiving element, thereby transferring a dye image to an image-receiving layer of said dye-receiving element to form said dye transfer image; and (b) thermally transferring a protection layer on top of said transferred dye image, said protection layer being applied from an element which contains inorganic particles, a polymeric binder and unexpanded synthetic thermoplastic polymeric microspheres, said microspheres having a particle size in the unexpanded condition of from 5 to 20  $\mu$ m, and which expand to 20 to 120  $\mu$ m upon application of heat during transfer of said protection layer to said image-receiving layer to provide a matte surface thereon, said microspheres comprising a mixture of low softening point microspheres and high softening point microspheres, said low softening point microspheres.

spheres having a softening point less than 105°C, said high softening point microspheres having softening point greater than 110°C, and the ratio of said low softening point microspheres to said high softening point microspheres being from 9:1 to 1:6.

## 5 10. A thermal dye transfer assemblage comprising

(a) a dye-donor element for thermal dye transfer comprising a support having thereon at least one dye layer area comprising an image dye in a binder and another area comprising a transferable protection layer, said transferable protection layer area being approximately equal in size to said dye layer area, wherein said transferable protection layer contains inorganic particles, a polymeric binder and unexpanded synthetic thermoplastic polymeric microspheres, said microspheres having a particle size in the unexpanded condition of from 5 to 20 μm, and which expand to 20 to 120 μm upon application of heat during transfer of said protection layer to a dye image-receiving layer of a dye-receiving element to provide a matte surface thereon, said microspheres comprising a mixture of low softening point microspheres and high softening point microspheres, said low softening point microspheres having a softening point less than 105°C, said high softening point microspheres to said high softening point microspheres being from 9:1 to 1:6; and

(b) a dye-receiving element comprising a support having thereon said dye image-receiving layer, said dye-receiving element being in a superposed relationship with said dye-donor element so that said dye layer is in contact with said dye image-receiving layer.



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# (54) Dye-donor element containing transferable protection overcoat

(57) A dye-donor element for thermal dye transfer comprising a support having thereon at least one dye layer area comprising an image dye in a binder and another area comprising a transferable protection layer, the transferable protection layer area being approximately equal in size to the dye layer area, wherein the transferable protection layer contains inorganic particles, a polymeric binder and unexpanded synthetic thermoplastic polymeric microspheres, the microspheres having a particle size in the unexpanded condition of from 5 to 20 μm, and which expand to 20 to 120 μm upon

application of heat during transfer of the protection layer to an image-receiving layer to provide a matte surface thereon, the microspheres comprising a mixture of low softening point microspheres and high softening point microspheres, the low softening point microspheres having a softening point less than 105° C, the high softening point microspheres having softening point greater than 110°C, and the ratio of the low softening point microspheres to the high softening point microspheres being from 9:1 to 1:6.



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